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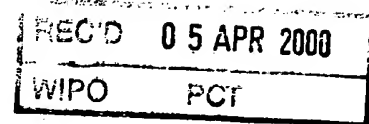
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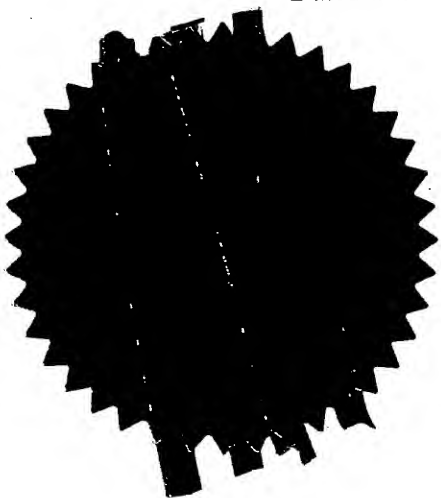
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I HEREBY CERTIFY that annexed hereto is a true copy of
documents filed in connection with the following patent
application:

Application No.	S990218
Date of Filing	18 March, 1999
Applicant	GAYA LIMITED, an Irish Company of 43 Fitzwilliam Place, Dublin 2, Ireland.

Dated this 18th day of March, 2000.

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PATENTS ACT, 1992

The Applicant(s) named herein hereby request(s)

- ☐ the grant of a patent under Part II of the Act
- ☒ the grant of a short-term patent under Part III of the Act

on the basis of the information furnished hereunder

1. **Applicant(s)**

Name: GAYA LIMITED

Address: 43 FITZWILLIAM PLACE
DUBLIN 2
IRELAND

Description/Nationality: An Irish Company

2. **Title of Invention:** A SURGICAL DEVICE

3. **Declaration of Priority on basis of previously filed application(s) for same invention (Sections 25 & 26)**

Previous Filing Date

Country in or for which Filed

Filing No.

4. **Identification of Inventor(s)**

Name(s) and Addresses of person(s) believed by Applicant(s) to be the Inventor(s)

Name(s):.....

Address:.....

5. Statement of right to be granted a patent (Section 2)(b))

6. Items accompanying this Request - tick as appropriate

- (i) ☒ prescribed filing fee (£50.00)
- (ii) ☐ specification containing a description and claims
☒ specification containing a description only
☒ drawings referred to in description or claims
- (iii) ☐ an abstract
- (iv) ☐ copy of previous application(s) whose priority is claimed
- (v) ☐ translation of previous application whose priority is claimed
- (vi) ☒ Authorisation of Agent (this may be given at 8 if this Request is signed by the Applicant(s))

7. Divisional Application(s)

The following information is applicable to the present application which is made under Section 24:-

Earlier Application No. Filing Date

Agent

The following is authorised to act as agent in all proceedings connection with the obtaining of a patent to which this request relates and in relation to any patent granted:-

MACLACHLAN & DONALDSON, 47 Merrion Square, Dublin 2

9. Address for Service (if different to that at 8)

MACLACHLAN & DONALDSON, at their address as recorded for the time being in the Register of Patent Agents (Rule 92)

Signed Name(s) GAYA LIMITED

By *A. M. McKearney*
MACLACHLAN & DONALDSON, Applicants' Agents

Date: 18th March 1999

A SURGICAL DEVICE

The present invention relates to a surgical device for use in minimally invasive surgery of the type using patient pneumoperitoneum and an access port.

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Minimally invasive surgery of this type is carried out having introduced gas into a patient's body cavity through an incision and sealed the incision with an access port. The access port enables laproscopic and hand or instrument assisted surgery to be performed.

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A sleeve forming such a port is shown in PCT Patent Application No. PCT/IE94/00045 entitled "Apparatus for use in surgery". The access port sleeve shown is used to create a controlled pressurized environment within the sleeve while allowing a surgeon's arm to pass through the sleeve. During surgery, gas is pumped into the patient's body cavity where the surgery is to be performed and the sleeve prevents gas escaping while allowing the surgeon to operate using minimally invasive surgery techniques. The application shows a sleeve having a flange at a distal end provided with adhesive for adhering the device to a patient's body or alternatively a mounting ring to surround the incision in a patient's body. While providing a suitable apparatus for performing such surgery the device described suffers from the principle disadvantage that in use, the sleeve protrudes upwardly from the patient and may interfere with the activities of the surgery team. Additionally, the sleeve must be sealed against the surgeon's upper forearm by clamping the device to the arm sufficiently tightly to avoid gas leak around the area of the seal. This presents the surgeon with a problem both in sealing the sleeve and in subsequent mobility.

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A further problem associated with the use of sleeves of the kind described is that a phenomenon known as "tenting" may occur. "Tenting" means that when the sleeve is adhered to the patient's skin or to a surgical drape and gas is induced into the patients abdominal cavity, there is a tendency for the sleeve to fill with gas and to pull away from the patient.

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There is therefore a need for a surgical device, which will overcome the aforementioned problems.

Accordingly, there is provided a surgical device for use in minimally invasive surgery of the type using an inflated body cavity accessible to a surgeon through an access port, defined by the device, surrounding an incision in a patients body, the device having: -

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body cavity engagement means for insertion into the incision to locate the device in position;

fixing means for attaching the device to a patients skin; and

sealing means connected between the body cavity engagement means and the fixing means, the sealing means being formed to prevent substantial leakage of gas from the body cavity on inflation when in an inoperative position and formed to mould to a substantial portion of a surgeon's hand or surgical instrument on

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insertion in an operating position.

Preferably, the body cavity engagement means is provided by an anchor ring formed for insertion into the incision.

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Preferably, the fixing means is provided by an adhesive web or fixing ring.

In one arrangement, the fixing means has an associated connector ring for receiving additional seals or medical instruments.

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Ideally, the sealing means is provided by a toroid cell formed to engage the incision between the fixing means and the body cavity engagement means.

Preferably, the cell forms a bladder through which the surgeon may access the body cavity, the bladder being filled with a viscous or semi-viscous liquid.

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Preferably, the bladder is filled with saline, gel or foam.

In one embodiment, the sealing means incorporates a foam shell formed for covering the incision.

5 Preferably, the foam shell is formed in two parts, or as a single part partially divided along one axis, the parts being movable relatively to allow a surgeon access to the body cavity.

In one arrangement the foam shell is formed by a plurality of individually disengageable layers. In this way the surgeon can adjust the height of the foam shell in response to particular needs by adding or removing foam layers. Thus a single device may be used on
0 abdomens of varying thickness, enhancing flexibility of application. Furthermore, the rigidity created by the induced gas and foam apron allows for hand insertion and withdrawal without the aid of an assistant or requiring the surgeon to use the other hand. Additionally, the external valve created by the inclusion of a foam shell is enhanced by the pressure of the induced gas passing up between the double walled tube and acting to force
15 the opposing faces of film together outside the patients abdominal cavity.

Preferably, the sealing means further incorporates a distal valve for insertion into the body cavity.

20 Ideally, the distal valve includes a mechanical seal.

The invention will now be described more particularly with reference to the accompanying drawings, which show, by way of example only, various embodiments of a surgical device in accordance with the invention, in which:-

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Fig. 1 is a top view of a surgical device in accordance with the invention;

Fig. 2 is a sectional view of the surgical device of Fig. 1 in the direction of the
arrows A-A;

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Fig. 3 is a sectional view similar to that shown in Fig. 2 showing the device in an inoperative position with a surgeon's hand approaching;

Fig. 4 is a sectional view as shown in Figs. 2 and 3 showing the device in an operating position with the surgeons hand in place;

5 Fig. 5 is a plan view of and alternative surgical device in accordance with the invention;

Fig. 6 is a front view of the surgical device of Fig. 5; and

Fig. 7 is an end view of the surgical device of Figs. 5 and 6.

Referring to the drawings, and initially to Figs. 1 to 4, there is illustrated a surgical device according to the invention, indicated generally by the reference numeral 1. The surgical device 1 is formed for use in minimally invasive surgery of the type using an inflated body cavity indicated generally by the reference numeral 2. The cavity 2 is accessible to a surgeon through an access port, defined by the device 1, surrounding an incision in a patient's abdominal wall 3.

15 In more detail, the device 1 has a body cavity engagement means provided by an anchor ring 5 for insertion into the incision to locate the device 1 in position. The device 1 is held in position on the patient's skin out side the body by a fixing means provided in this case by an adhesive web 6. The ring 5 and web 6 ensure that the device 1 is securely fixed in position and surround the incision. It will be noted that the web may be replaced by any functional equivalent to secure the device in position.

25 The web 6 has an associated connector ring 7 for receiving additional seals to prevent loss of pressure from the cavity 2. The connector ring 7 may also be used for holding or guiding medical instruments into position over or in the incision.

30 The device 1 has a sealing means, provided in this embodiment of the invention, by a saline filled toroid cell 8 connected between the anchor ring 5 and the web 6. The cell 8 is formed to prevent substantial leakage of gas from the body cavity 2 on inflation when in an

inoperative position see Figs. 2 and 3. The cell 8 is also formed to mould to a substantial portion of a surgeon's hand or surgical instrument when in an operating position (see Fig. 4). The cell 8 is also formed to allow for the removal of operative tissue when in an operating position with or without pneumoperitoneum established.

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It will be noted that the cell may be filled with any suitable material and represents a significant improvement over prior art devices, which are inflated with air. The use of a liquid such as a sealed saline bladder improves hygiene around the wound and responds more quickly to a movement by a surgeon's hand. Additionally the invention overcomes problems associated with inflatable bladders, which will leak air if under inflated or be overly restrictive to movement if over inflated.

It will further be noted that the sealing means is described as a toroid or donut shaped cell, but that it could be equally provided as a lip shaped or elliptical cell tapering slightly at either end.

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In use, an incision is made in the abdominal wall 3 and the anchor ring 5 passed through the incision into the cavity 2. The anchor ring 5 is moved when in the cavity 2 such that the ring 5 surrounds the incision. The web 6 is then attached to the patients skin to fix the device 1 in position with the cell 8 being connected between the web 6 and the ring 5 and engaging the portions of the abdominal wall 3 exposed by the incision. The cell 8 seal the incision and the abdominal cavity 2 may be inflated as required by the surgeon to an inoperative position Fig.2. The surgeon can gain access to the cavity 2 losing a minimum of gas pressure by passing a hand or instrument through the center of the toroid or donut shaped cell 8. When the hand or instrument is in the operating position (Fig. 4) the cell moulds to the hand or instrument to prevent loss of pressure.

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Referring now to Figs. 5 to 7 there is illustrated a further surgical device in accordance with the invention indicated generally by the reference numeral 20, in which parts similar to those identified with reference to Figs. 1 to 4 are identified by the same reference numerals generally. In this embodiment the sealing means is in two sections. A foam shell 28 is in this case formed in two parts to envelop the incision site. It will be understood that

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the foam shell may equally be provided as a single part, divided or split along an axis. The parts of the shell 28 are movable relatively to allow a surgeon access to the body cavity and can be biased together to seal the cavity 2 when not in use. A sleeve 30 connected to the web 6 covers the shell 28 and passes into the cavity 2 and is terminated in the cavity 2 by a distal valve 31 having a mechanical seal 32. It will also be understood that the web 6 may equally be provided by an anchoring ring.

In use, the parts of the foam shell 28 are separated as before by the surgeon's hand or instrument and the foam moulds the shape of the inserted object to prevent loss of pressure. The inserted object then travels through the sleeve 30 to the distal valve 31 inside the cavity 2 and opens the mechanical seal 32. When the task has been completed the inserted object is removed and the mechanical seal 32 being so biased closes. The pressure in the cavity 2 is such during procedures of this type to close the sleeve 30 along its length as the object is removed and a final seal is provided by the foam shell 28 decompressing when the object has been removed.

The use of a foam shell has a number of advantages over known systems. For example, trauma at the incision is minimised as shock associated with downward pressure when inserting the surgeon's hand is largely absorbed by the foam. Tenting is eliminated as the foam shell reduces the volume of gas in the proximal end of the sleeve. The foam may also be used to absorb liquids such as blood in a hygienic manner and may reduce the effect of blood and body fluids on the anchoring ring. Furthermore it is envisaged that the lifting action of the foam may be used to retract tissue or for creating additional anchoring forces or between the distal valve and the abdominal wall. The cell may also be formed in any suitable manner to allow for the removal of operative tissue during the course of an operation whether or not pneumoperitoneum has been established.

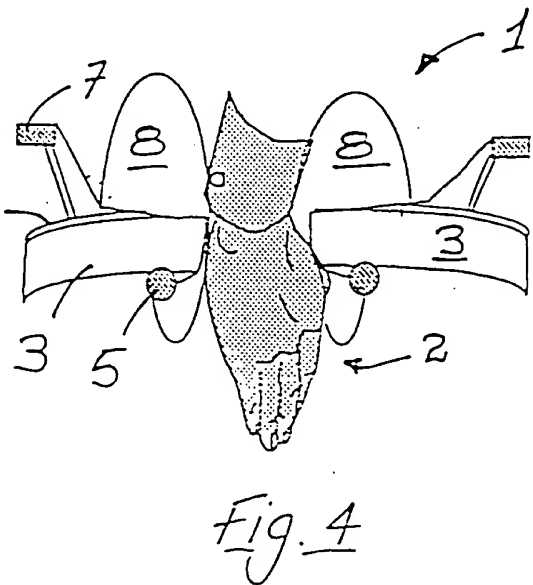
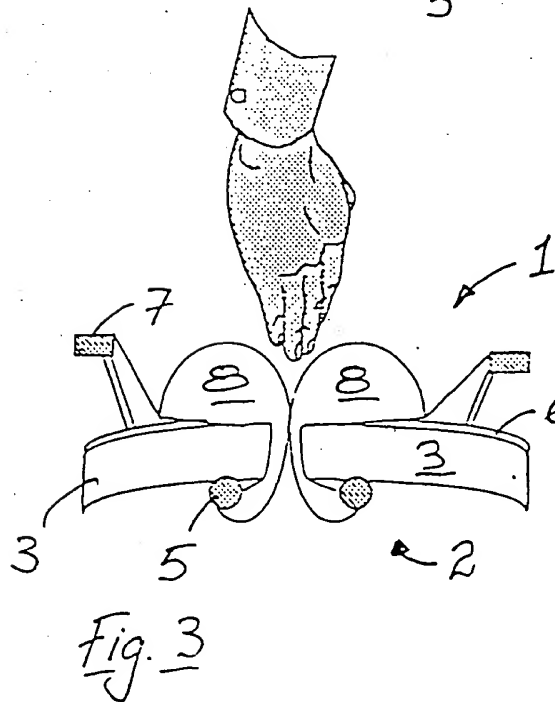
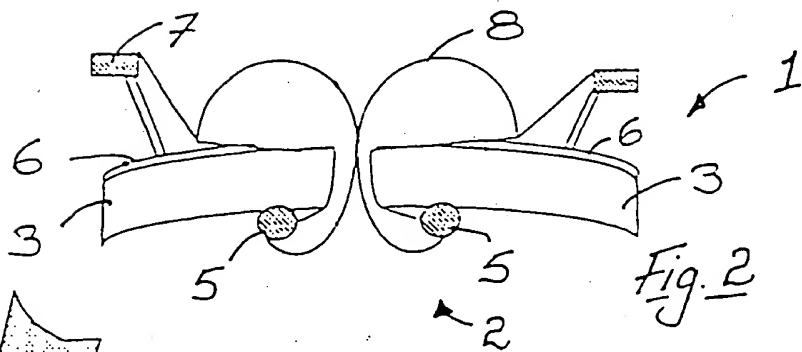
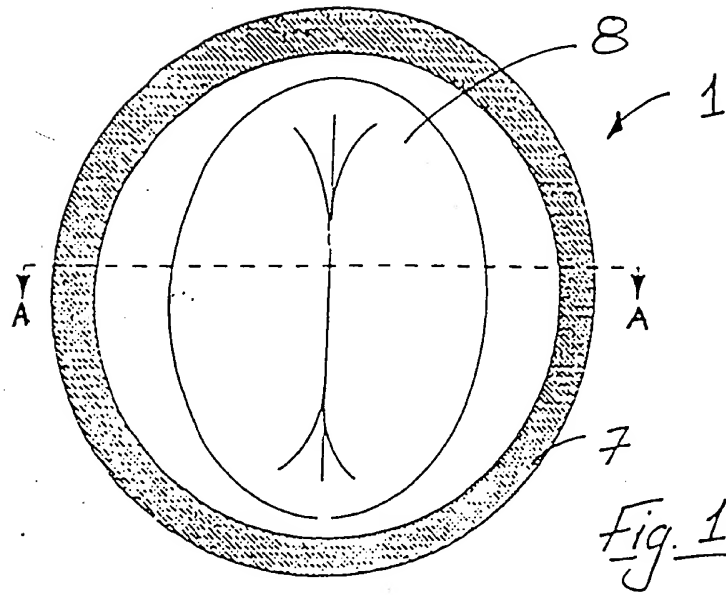
It will be noted that the sleeve and valve may incorporate means for preventing the sleeve returning through the incision accidentally. These means may include but are not limited to an angled flap or flaps on the distal valve, tensioning means in the sleeve such as weld lines or a physical connection to a body part including the abdominal wall.

It will be understood that the foam shell may also be provided as a single block, defining a passageway therein, to allow communication between the exterior and the cavity.

5 It will of course be understood that the invention is not limited to the specific details described herein, which are given by way of example only, and that various modifications and alterations are possible within the scope of the invention.

MACLACHLAN & DONALDSON,
Applicant's Agents,
47, Merrion Square,
DUBLIN 2.

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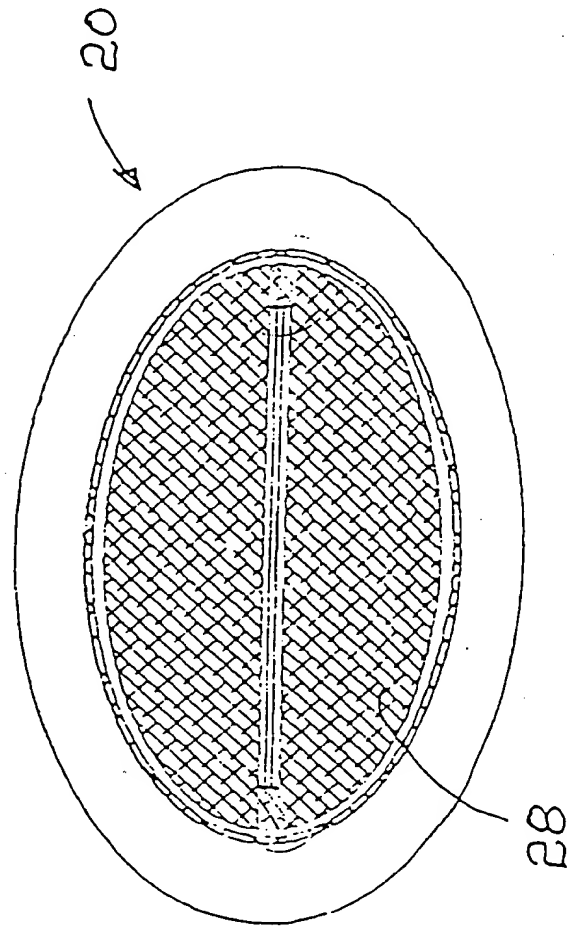
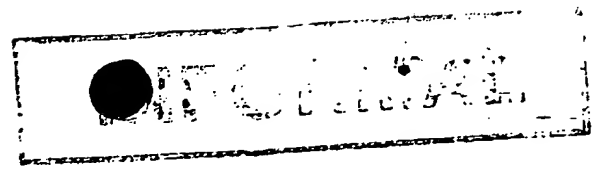


Fig. 5

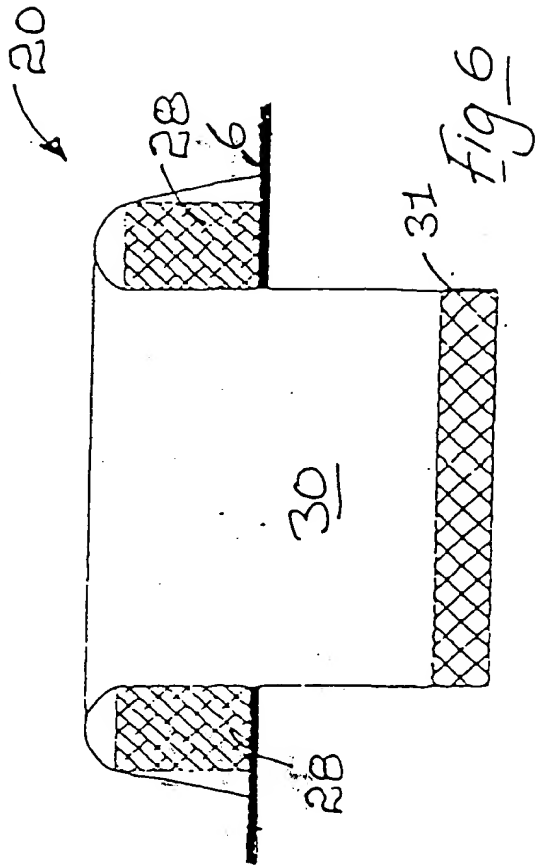


Fig. 6

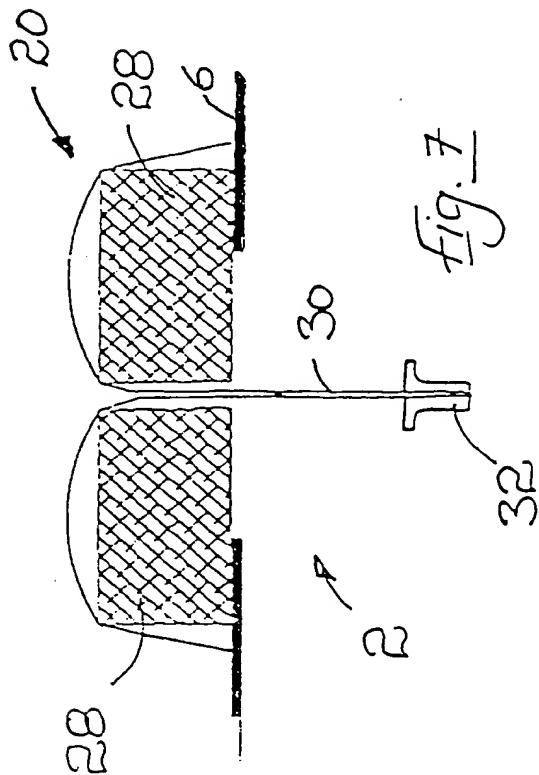


Fig. 7